

1 **Title:** The Roles of Drift and Selection on Short Stamen Loss in *Arabidopsis thaliana* along an Elevational
2 Gradient in the Spanish Pyrenees

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46

47 **Abstract**

48 Traits that have lost function sometimes persist through evolutionary time. These traits may be
49 maintained by a lack of standing genetic variation for the trait, if selection against the trait is weak
50 relative to drift, or if they have a residual function. To determine the evolutionary processes shaping
51 whether nonfunctional traits are retained or lost, we investigated short stamens in 16 populations of
52 *Arabidopsis thaliana* along an elevational cline in the Spanish Pyrenees. We found a cline in short
53 stamen number from retention of short stamens in high elevation populations to incomplete loss in low
54 elevation populations. We did not find evidence that limited genetic variation constrains the loss of
55 short stamens at high elevations nor evidence for divergent selection on short stamens between high
56 and low elevations. Finally, we identified loci associated with short stamens in the Spanish Pyrenees that
57 are different from loci associated with variation in short stamen number across latitudes from a
58 previous study. Overall, we did not identify the evolutionary mechanisms maintaining an elevational
59 cline in short stamen number but did identify different genetic loci underlying the variation in short
60 stamen along similar phenotypic clines.

61 **Keywords (provide 3 to 6):** – genetic drift, trait loss, selection, effective population size, stamen,
62 elevational cline

63 **Teaser text**

64 The evolutionary mechanisms underlying loss or retention of traits that have lost function are poorly
65 understood. Short stamens in *Arabidopsis thaliana* provide a compelling system to investigate the roles
66 of genetic drift and selection in trait loss across latitudinal and elevational clines. This study investigates
67 the role of drift and selection in short stamen loss in 16 populations of *A. thaliana* along an elevational
68 gradient in the Spanish Pyrenees. An investigation of the genetic loci underlying variation in short
69 stamen number suggests mutations in different genes may cause trait loss in similar phenotypic clines
70 within a species.

71

72 Introduction

73 Traits that have lost function are often lost through evolutionary time, yet some persist. Nonfunctional
74 traits could be lost through direct selection against the trait (Dorken et al., 2004; Lahti et al., 2009),
75 correlated responses to selection on other traits caused by pleiotropy or linkage disequilibrium
76 (Yoshizawa et al., 2012), or the accumulation of selectively neutral mutations (Fong et al., 1995). In
77 contrast, nonfunctional traits may be maintained by evolutionary constraint if the nonfunctional trait is
78 genetically correlated with a different, functional trait (Lande, 1979; Walsh & Blows, 2009).
79 Nonfunctional trait loss could also be prevented if there is not enough standing genetic variation for
80 selection to act on the trait (Lahti et al., 2009) or if weak selection against the trait is unable to
81 overcome drift (Charlesworth, 2009). Both of these processes may be pronounced in species with small
82 effective population sizes. Determining the evolutionary processes shaping whether nonfunctional traits
83 are retained or lost will have broad implications for our understanding of the interplay of direct
84 selection, correlated responses to selection, and genetic drift (reviewed in Futuyma, 2010).

85 Here, we use an elevational cline in short stamen number in *Arabidopsis thaliana* to investigate how
86 evolutionary processes interact to shape variation in a trait that has lost function. Almost all of the 3,700
87 species in the Brassicaceae family have flowers with four long and two short stamens (Endress, 1992).
88 While the function of short stamens is unknown, maintaining the stamen length difference is likely
89 adaptive in wild radish, *Raphanus raphanistrum*, an outcrossing species within Brassicaceae (Conner et
90 al., 2003; Waterman et al., 2023). However, *A. thaliana* evolved to be almost entirely self-fertilizing from
91 an outcrossing ancestor between 0.5 – 1 million years ago (Durvasula et al., 2017; Tang et al., 2007). In
92 flowering plants, a transition from outcrossing to self-pollination and subsequent relaxation of selection
93 for pollination often results in a suite of trait changes, called the ‘selfing syndrome’, that includes
94 smaller flowers and reduced distance between the anthers and stigma (Sicard & Lenhard, 2011). Self-
95 pollination also decreases local effective population size (Caballero, 1994), increasing the effects of
96 genetic drift.

97 Short stamens in *A. thaliana* do not contribute significantly to self-fertilized seed number, but have
98 nonetheless been retained in most natural populations (Royer et al., 2016). In particular, southern
99 European populations are more likely to lose short stamens while northern European populations retain
100 both short stamens. Because northern European populations of *A. thaliana* have undergone repeated
101 bottlenecks that reduced effective population size, and thus standing genetic variation (Beck et al.,
102 2008; François et al., 2008; Lewandowska-Sabat et al., 2010), short stamens may be retained in these
103 populations because there is insufficient genetic variation for stamen number and/or weak selection
104 against stamens cannot overcome drift. Alternatively, the latitudinal cline of the European populations
105 could be maintained by local adaptation for short stamen number or another correlated trait. The same
106 evolutionary processes may shape elevational clines in short stamen number. High elevations were
107 glaciated, even at southern latitudes (Hughes et al., 2006) and while the Spanish Pyrenees do include
108 putative refugia regions (Médail & Diadema, 2009), prior work in the Pyrenees has shown high elevation
109 montane populations have less genetic variation and smaller effective population sizes than lower
110 elevation coastal populations (Gomaa et al., 2011; Montesinos et al., 2009). Further, because
111 environmental factors vary similarly with latitude and elevation, selection may result in similar
112 phenotypic clines. Yet, latitudinal and elevational phenotypic clines are not always parallel (Daco et al.,
113 2021; Kooyers et al., 2015) and when parallel phenotypic clines are identified, they may not have a
114 parallel genetic basis (Fulgione et al., 2022; Gamba et al., 2022).

115 To understand the processes shaping variation in short stamen loss, we assessed 16 populations of *A.*
116 *thaliana* along an elevational gradient in the Spanish Pyrenees. We counted short stamen number in a
117 growth-chamber common garden and sequenced multiple individuals per population to quantify genetic
118 variation. We identified an elevational cline in short stamen number in the Spanish Pyrenees, with more
119 stamen loss at low elevation, similar to the previously described latitudinal cline (Royer et al., 2016). We
120 then used multiple regression, polygenic adaptation detection, and genome wide association studies to
121 address possible mechanisms for maintaining the cline:

- 122 **1)** Is the short stamen loss cline maintained by variation in effective population size? Less
123 stamen loss in populations with less genetic variation after accounting for variation due to
124 elevation would be consistent with the hypothesis that stamen loss is constrained by a lack
125 of genetic variation and less effective selection.
- 126 **2)** Is the short stamen loss cline maintained by local adaptation? Evidence for divergent
127 selection on short stamen number between high and low elevation populations would
128 support local adaptation maintaining the cline.
- 129 **3)** What genomic loci are associated with short stamen number? If we identify loci within the
130 three previously identified QTL (Royer et al., 2016), then this would support parallel genomic
131 evolution between the latitudinal and elevational clines in short stamen number.

132 **Materials and Methods**

133 *Seed Collection*

134 Population collection sites are along an elevation gradient across the Pyrenees and along the
135 Mediterranean coast (Fig. 1A, Table S1). Seeds from five to nine individuals were haphazardly collected
136 from multiple patches in each of 16 populations ($n = 112$ genotypes) in northeast Spain (Montesinos-
137 Navarro et al., 2011). These lineages are also used in Montesinos-Navarro et al. (2011), Montesinos-
138 Navarro et al. (2012), and Wolfe and Tonsor (2014). They include the 10 populations from Montesinos et
139 al. (2009) and the 9 populations from Gomaa et al. (2011).

140 *Phenotyping*

141 To assess genetic differentiation for short stamen number, plants were grown in chamber common
142 gardens in three blocks at Michigan State University, East Lansing, MI. Seeds were stratified at 4°C for
143 five days before we increased temperatures to 22°C:18°C for 16-hour days at a constant 60% humidity
144 for four weeks. After emergence, plants were thinned to one seedling per cell in a 200 cell plug tray.
145 One to four plants from each of the 112 genotypes (median = 2, $n = 230$ plants) were vernalized at 4°C
146 with 10-hour days for six weeks before returning to 22°C:18°C for 16-hour days through flowering.

147 On each plant, we counted stamen number (Fig. 1B) on up to three flowers at each of up to three
148 timepoints throughout the duration of flowering to estimate short stamen number (1 to 9 flowers per
149 plant, median = 6, $n = 1,423$ flowers). Population mean short stamen number was calculated as the
150 arithmetic mean from all flowers phenotyped in a population (42 to 169 flowers per population, mean =
151 88.9, median = 99.5). The arithmetic population means were highly correlated with estimated marginal
152 means for each population calculated with the *emmeans* package (Lenth, 2021) in R (v4.2.2 (R Core
153 Team, 2021)) from a model with population as a fixed effect, random effects of flowering timepoint

154 nested within plant nested within genotype which is nested within population, and random effect for
155 block ($r = 0.975$, $p < 0.001$).

156 We sequenced the entire genomes of a subset of 61 genotypes (see below). The 61 genotypes include
157 representatives from all 16 populations (3 or 4 different genotypes per population). In the sequenced
158 genotypes, stamen number was counted on up to four plants per genotype (median = 2, $n = 141$) and up
159 to nine flowers per plant (median = 7, $n = 971$). For all analyses that incorporate both phenotypic and
160 SNP information, arithmetic population means were recalculated as the mean from flowers scored for
161 short stamen number from sequenced genotypes (18 to 118 flowers per population, median = 63). The
162 population means from all flowers and the population means from only the sequenced genotypes are
163 highly correlated ($r = 0.985$, $p < 0.001$) indicating the sequenced genotypes are representative of all
164 plants scored for short stamen number.

165 *Sequencing*

166 A subset of 61 genotypes (3-4 per population) were chosen for Illumina paired-end whole-genome 150bp
167 sequencing using the WGS-Novogene platform. Nextera adapter sequences were trimmed from the raw
168 sequence data with Trim Galore (Krueger, 2019). We also clipped the first 15 bp of each read because
169 quality checks with FastQC (Andrews, 2019) and MultiQC (Ewels et al., 2016) showed an identical,
170 unusual, pattern in the first 10 bases of each read: GTTTTAAACT. Reads were mapped to the TAIR10
171 reference genome (Berardini et al., 2015) using BWA mem with default settings (Li & Durbin, 2009). The
172 mean mapping rate for properly paired reads was 96.6% with a mean of 15,224,790 properly paired and
173 mapped reads per genotype and a total of 943,936,960 mapped and paired reads in the dataset. The
174 median depth across genotypes was low but acceptable at 8X. There is variation in coverage, missing
175 data, and quality scores between genotypes, but no genotypes were excluded on this basis (Table S2).
176 Duplicate reads were marked with the Genome Analysis Toolkit (GATK) v4.1.4.1 (Van der Auwera &
177 O'Connor, 2020) MarkDuplicates Spark. After an initial round of Haplocaller and GenotypeGVCF, the
178 dataset was filtered with the parameters suggested by GATK best practices (Van der Auwera &
179 O'Connor, 2020): $QD < 2$, $FS > 60$, $MQ < 40$, $ReadPosRankSum < -8$, and $MQRankSum < -12.5$. The filtered
180 file was used as a known variants file for base quality score recalibration (BQSR). Variants were then
181 called again with HaplotypeCaller using the GVCF flag to keep all sites before combining samples for
182 GenotypeGVCF with the all-sites flag to create a dataset that includes both variant and invariant sites
183 (116,855,685 total sites).

184 *Testing the drift hypotheses*

185 To estimate the effects of past drift and gene flow in each population, genome-wide pairwise nucleotide
186 diversity (π) was calculated for each population as an approximate measure of effective population size
187 with pixy v1.0.4 (Korunes & Samuk, 2021) from a filtered dataset containing variant and invariant sites
188 based on the pixy protocol. We used quality filters to remove all indels and low-quality sites that met
189 the following criteria: quality score less than 20, mean depth less than 3, and more than 25% missing
190 data. The filtered dataset contains 99,202,614 sites. A peak in nucleotide diversity was observed in the
191 centromere of each chromosome, likely caused by fewer mapped sites (Korunes and Samuk, pers. com.).
192 However, excluding previously published centromeres (Clark et al., 2007) did not meaningfully change
193 genome-wide nucleotide diversity ($r = 0.998$, $p < 0.001$), so centromeres are included in all of the
194 following analyses (results excluding the centromeres can be found in Figs. S5-S7).

195 To test if variation in effective population size is maintaining a cline in short stamen number, we used
196 multiple regression to identify how short stamen number was predicted by nucleotide diversity and
197 elevation. While elevation is correlated with climatic variables such as precipitation and temperature
198 (Montesinos-Navarro et al., 2011) that could be selective agents for plant traits, we used elevation
199 because previous work in these same 16 populations demonstrated that elevation explained 54% of the
200 variance in trait principal component 1 (traits include: phenology, water use efficiency, instantaneous
201 CO₂ and H₂O exchange, PSII quantum efficiency, and specific leaf area) while the climate PC1 only
202 explained 36% of the trait variation (Wolfe & Tonsor, 2014). We then visualized these results by using
203 the residuals of single regression models. We used *lme4* in R for all models (Bates et al., 2015).

204 A negative relationship between nucleotide diversity and short stamen number after correcting for
205 elevation would support the hypothesis that smaller effective population sizes are constraining trait loss
206 due to low standing variation and less effective selection. A relationship between short stamen number
207 and elevation after correcting for nucleotide diversity could be evidence for divergent selection because
208 it indicates the short stamen number cline persists beyond the variation explained by nucleotide
209 diversity. However, this relationship could also result from shared evolutionary history, sometimes
210 called shared genetic drift, if relatedness among populations is associated with elevation (Colautti & Lau,
211 2015). Therefore, we further investigated evidence for divergent selection by testing for local adaptation
212 in short stamen number.

213 *Testing for local adaptation*

214 We used plink v1.9 (Chang et al., 2015) to filter the all-sites output from GATK to a “variant sites only”
215 dataset for conducting genetic principal component analysis (PCA). The “variant sites only” dataset was
216 also used for a genome-wide association study (GWAS; see below). We used quality filters to remove all
217 indels and non-biallelic sites in addition to low quality sites that met the following criteria: minor allele
218 frequency less than 5%, quality score less than 25, mean depth less than 5, and more than 25% missing
219 data. The variant sites dataset contains 1,858,706 SNPs. Filtering parameters were chosen to maintain
220 the same percent nonsynonymous sites (annotated by snpEff (Cingolani et al., 2012)) as stricter
221 parameters while retaining more variants in the dataset. Genetic PCA was calculated for the first 20 PCs
222 with plink v1.9 (Chang et al., 2015) to characterize genetic relatedness within, and differentiation
223 between, populations. We looked for correlations between the population average PC value and
224 elevation for the first 4 PCs to identify population structure associated with elevation.

225 We tested for divergent selection across populations on short stamen number, i.e., selection for more
226 stamens at high elevation and/or fewer stamens at low elevation than expected due to genetic drift,
227 with Qpc using the *quaint* R package (Josephs et al., 2019). Qpc is an extension of Qst-Fst analysis that
228 uses genetic PCs to estimate additive genetic variance within and between populations (Josephs et al.,
229 2019), and tests for phenotypic differentiation due to selection beyond that expected from neutral
230 evolution. This differs from the multiple regression test for the drift hypothesis described above because
231 Qpc explicitly considers among population differentiation. Qpc incorporates genetic relatedness within
232 and among populations through a kinship matrix, rather than a single measure of population
233 differentiation as in Qst-Fst. Qpc then tests for excess phenotypic divergence along major axes of
234 genetic relatedness (principal components of the kinship matrix) rather than excess phenotypic
235 divergence between populations as in Qst-Fst (Josephs et al., 2019). The input kinship matrix was
236 generated from a random subset of 50,000 SNPs that had no missing data. This kinship matrix is slightly

237 different from the kinship matrix used for genetic PCA because Qpc requires a kinship matrix
238 standardized across all loci, not each locus individually as in plink, and *quaint* is not capable of dealing
239 with missing data.

240 *Using genome wide association studies (GWAS) to identify loci associated with short stamen loss*

241 We performed GWAS to find genomic regions associated with short stamen number. Mean short
242 stamen number was not normally distributed (Shapiro-Wilk $w=0.85$, $p = 3.27 \times 10^{-6}$; Fig. S1). Arcsine
243 transformation made the distribution closer to normal but still skewed (Shapiro-Wilk $w = 0.91$, $p =$
244 2.72×10^{-4} ; Fig. S1). Phenotypic and genotypic data were merged in plink v1.9 (Chang et al., 2015). GWAS
245 was conducted in gemma v0.98.4 with the Wald hypothesis test for each SNP against the alternate allele
246 and a centered kinship matrix to account for population structure (Zhou & Stephens, 2012). The output
247 was visualized with the *qqman* (Turner, 2018) and *ggplot2* (Wickham et al., 2019) R packages. SNPs were
248 further investigated if they passed a significance threshold specified using a false discovery rate (FDR) <
249 0.05 determined with *p.adjust* in R.

250 Additional GWAS were conducted due to the remaining skew in the arcsine-transformed data. We ran a
251 GWAS with untransformed mean short stamen number and another with mean short stamen number
252 coded as a binary trait. In the latter, genotypes with no short stamen loss (mean short stamen number =
253 2) were coded as controls and genotypes with any amount of short stamen loss (mean short stamen
254 number < 2) were coded as cases. Finally, a fourth GWAS was conducted on the subset of genotypes
255 that experience short stamen loss (i.e., only the genotypes with mean short stamen number < 2;
256 Shapiro-Wilk $w=0.93$, $p=0.0132$) to ameliorate the zero-inflated-like distribution in the other continuous
257 GWAS caused by excess genotypes with a mean short stamen number equal to 2 (Fig. S1).

258 We identified genetic regions with SNPs associated with short stamen number in at least two GWAS to
259 find candidate regions for further study (“shared” SNPs). We identified overlapping regions associated
260 with stamen loss among the four GWAS analyses by creating a 1kb window centered on each SNP that
261 had an FDR adjusted p-value below 0.10 in any GWAS and searching for SNPs within that window that
262 were also below an FDR adjusted p-value of 0.10 in at least one other GWAS. We chose a more lenient
263 FDR of 0.10 for this analysis because we have higher confidence SNPs are associated with short stamen
264 number if they pass a significance threshold in multiple analyses. We chose a window of 1kb to include
265 regions flanking the associated SNP, although we recognize that *A. thaliana* genes can be larger than 1kb
266 and that linkage disequilibrium (LD) begins to drop off around 50kb in our individuals (Fig. S2), so this is
267 a conservative overlap criterion. SNPs in overlapping stamen loss associated regions were investigated
268 with the TAIR10 genome browser.

269 All figures were created with *ggplot2* (Wickham et al., 2019) and *ggpubr* (Kassambara, 2023) unless
270 otherwise noted.

271 **Results**

272 *Short stamen loss is more common at low elevation*

273 Mean short stamen number, when measured in a common garden, increases with the elevation of the
274 source population until reaching close to two stamens around 1300m ($\beta = 4.84 \times 10^{-4}$, $p = 0.004$; $\gamma = -$
275 4.73×10^{-7} , $p = 0.137$; Fig. 2A; Table S3). Thus, short stamen retention is more common in populations
276 from higher elevations and loss is more common at low elevations in the Spanish Pyrenees. This

277 elevational cline is similar to the previously identified latitudinal cline where short stamen loss is more
278 common at southern latitudes (Royer et al., 2016).

279 *Nucleotide diversity decreases with elevation, but there is no evidence that variation in effective*
280 *population size maintains the cline in short stamen number*

281 Consistent with expectations that *A. thaliana* in the Pyrenees experienced repeated founder effects
282 after the last glaciation, high elevation populations have less nucleotide diversity than low elevation
283 populations ($\beta = -1.81 \times 10^{-6}$, $p < 0.001$; Fig. 2B; Table S3). The three populations with the lowest
284 nucleotide diversity are high elevation populations BIS, VIE, and PAN. The results are consistent with
285 prior findings that genetic diversity and effective population size decrease with elevation in this region
286 (Gomaa et al., 2011; Montesinos et al., 2009). Population nucleotide diversity is comparable to
287 nucleotide diversity of *A. thaliana* across the Iberian Peninsula and the European range (Alonso-Blanco
288 et al., 2016).

289 We hypothesized a negative relationship between nucleotide diversity and short stamen number if
290 variation in effective population size was maintaining the short stamen cline, but nucleotide diversity did
291 not significantly predict short stamen number when accounting for elevation ($\beta = -138.5$, $p = 0.167$; Fig.
292 2D; Table S3). Further, the relationship between short stamen number and elevation is similar whether
293 nucleotide diversity is included or not (Fig. 2C, compare to Fig. 2A; Table S3). These results suggest the
294 short stamen number cline is not maintained by variation in effective population size but may be
295 maintained by local adaptation, correlated responses to selection, or shared evolutionary history.

296 *There is no evidence for divergent selection on short stamen number along the elevational cline*

297 An elevational cline in short stamen number could result from local adaptation to different optima at
298 high and low elevations. To demonstrate evidence of local adaptation resulting from divergent selection,
299 we would need to show that the cline in short stamen number is stronger than what could be generated
300 by neutral evolution alone. Genetic structure, here measured using genetic PCA, shows a strong pattern
301 of genetic differentiation by elevation; both PC1 ($r = -0.75$, $p = 9.22 \times 10^{-4}$) and PC2 ($r = -0.54$, $p = 0.03$) are
302 correlated with elevation (Fig. 3A). Neither PC3 nor PC4 are correlated with elevation (Fig. S3). The BOS
303 population is an outlier along both PC1 and PC2 (Fig. 3A). This aligns with geographic information
304 because BOS is located on a northern face of the Spanish Pyrenees while the other populations are on
305 southern faces, and prior work identified BOS in the Northwestern genetic cluster of the Iberian
306 Peninsula while the other 15 populations belong to the Northeast cluster or are classified as mixed
307 (Castilla et al., 2020). The PCA was conducted a second time after removing the BOS population (Fig. 3B).
308 These results continue to show an elevational cline along PC1 ($r = -0.92$, $p = 1.15 \times 10^{-6}$), though PC2 ($r = -$
309 0.17 , $p = 0.55$) separates the four highest elevation populations from each other with the closest
310 clustering of PAN and VIE. The strong genetic differentiation across elevation we observed means that
311 we have low power to test for even greater differentiation in stamen loss as evidence for divergent
312 selection. Not surprisingly, we did not find evidence for divergent selection on short stamen number
313 with *Qpc*. Differences in short stamen number between high and low elevation populations were not
314 larger than could be explained by neutral evolution (Fig. 3C, $p = 0.257$).

315 *Few SNPs across the genome are associated with short stamen number*

316 We conducted GWAS to identify loci associated with short stamen number. Because short stamen
317 number was not normally distributed (Fig. S1), we conducted GWAS on arcsine-transformed mean short
318 stamen number, untransformed mean short stamen number, a binary trait of whether or not individuals
319 lacked any short stamens, and an untransformed subset of only genotypes that lack some short
320 stamens. Each GWAS found different SNPs associated with the trait. The qqplot for the arcsine-
321 transformed GWAS shows most p-values close to the 1:1 line between Expected and Observed (Fig. 4)
322 but the p-values in the binary GWAS deviate early from the 1:1 line, indicating a high false positive rate
323 in identifying SNPs associated with any short stamen loss (Fig. S4). In accordance with this observation,
324 the binary GWAS had 1,707 SNPs associated with short stamen loss at FDR < 0.05 while the other
325 analyses only had two or three (Figs. 4, S4). Only two SNPs are associated with arcsine-transformed
326 short stamen number (FDR < 0.05; Fig. 4). The SNP at Chr3:2942726 is within a FAD/NAD(P)-binding
327 oxidoreductase family protein (AT3G09580) that localizes in the chloroplast (Tomizioli et al., 2014). The
328 SNP at Chr5:13458838 is not within a gene; the closest gene, PICALM3 (AT5G35200), is just over 3kb
329 away.

330 We identified top candidates for loci associated with short stamen number by identifying 1kb windows
331 that have SNPs associated with short stamen number in multiple GWAS (“shared” SNPs). To do so, we
332 calculated 1kb windows centered on all SNPs associated with short stamen number (FDR < 0.10) and
333 searched for other SNPs that fit the same criteria and fell within each window. Twenty SNPs were
334 associated with short stamen number in more than one GWAS from 9 different 1kb windows (Table S4).
335 None of the SNPs fall within genes with known stamen function. However, a shared SNP at
336 Chr3:2253161 is approximately 40kb away, thus within LD (Fig. S2), from *FHA2* (AT3G07220), a
337 SMAD/FHA domain containing protein involved in stamen development (Ahn et al., 2013; Gu et al.,
338 2020). Mutating *FHA2* proteins can cause plants to have fewer stamens, though flowers sometimes lose
339 long stamens and sometimes lose short stamens (Ahn et al., 2013). The two SNPs associated with
340 arcsine-transformed short stamen number (FDR < 0.05) are included in the shared SNPs (Fig. 4; Table
341 S4).

342 All of the SNPs associated with variation in short stamen number fall outside the 95% credible intervals
343 of previously identified QTL for stamen loss using recombinant inbred lines with parents from the
344 extremes of the latitudinal stamen loss gradient (Royer et al., 2016). The closest intersection is on
345 chromosome 5, where SNPs associated with short stamens are approximately 2,000kb further into the
346 chromosome than the previously identified stamen loss QTL. This is beyond the start of LD decay we
347 estimated of 50kb (Fig. S2).

348 Discussion

349 In this study, we tested for an elevation cline in short stamen loss. We found that short stamen number
350 increased with elevation (Fig. 2A), similar to the latitudinal cline observed by Royer et al. (2016). Short
351 stamen number is one of many traits that show an elevational cline in this region (Montesinos et al.,
352 2009; Montesinos-Navarro et al., 2011; Wolfe & Tonsor, 2014). Of these traits, days to bolting shows
353 similar elevational and latitudinal clines with delayed bolting at high elevations and northern latitudes
354 (Montesinos-Navarro et al., 2011; Stinchcombe et al., 2004). Thus, short stamen number joins a growing
355 database of traits with an elevational cline in the Spanish Pyrenees and a much smaller set of traits with
356 parallel elevational and latitudinal clines. These clines could be maintained by effective population sizes
357 because both high elevation populations and northern latitude populations may have undergone

358 repeated bottlenecks that reduced effective population size since the last glaciation (Beck et al., 2008;
359 François et al., 2008; Hughes et al., 2006; Lewandowska-Sabat et al., 2010). While the Pyrenees do
360 include putative glacial refugia regions (Médail & Diadema, 2009), none of the 16 populations analyzed
361 here have been classified as relict lineages that predate the last glaciation (Castilla et al., 2020).
362 Selection could also maintain similar latitudinal and elevational clines because environmental factors
363 such as temperature often vary in the same way across latitude and elevation.

364 We then tested if the cline in short stamen loss was maintained by variation in effective
365 population size if a combination of low genetic variation and less effective selection in high
366 elevation populations hinders a response to selection against short stamens. We found
367 variation in genetic diversity across elevations consistent with expectations from repeated
368 founder effects during range expansion after the last glaciation and/or low gene flow (Fig.
369 2B). However, the multiple regression suggests that variation in genetic diversity is not
370 maintaining the short stamen number cline (Fig. 2C and 2D). Instead, the multiple regression
371 suggests that elevation can explain both variation in genetic diversity and in stamen number.
372 It is important to note that our clines show populations with low genetic diversity maintain
373 two stamens. This is opposite to the hypothesis that genetic drift would cause trait loss which
374 has been shown in other systems (Eckert et al., 1996; Lahti et al., 2009).

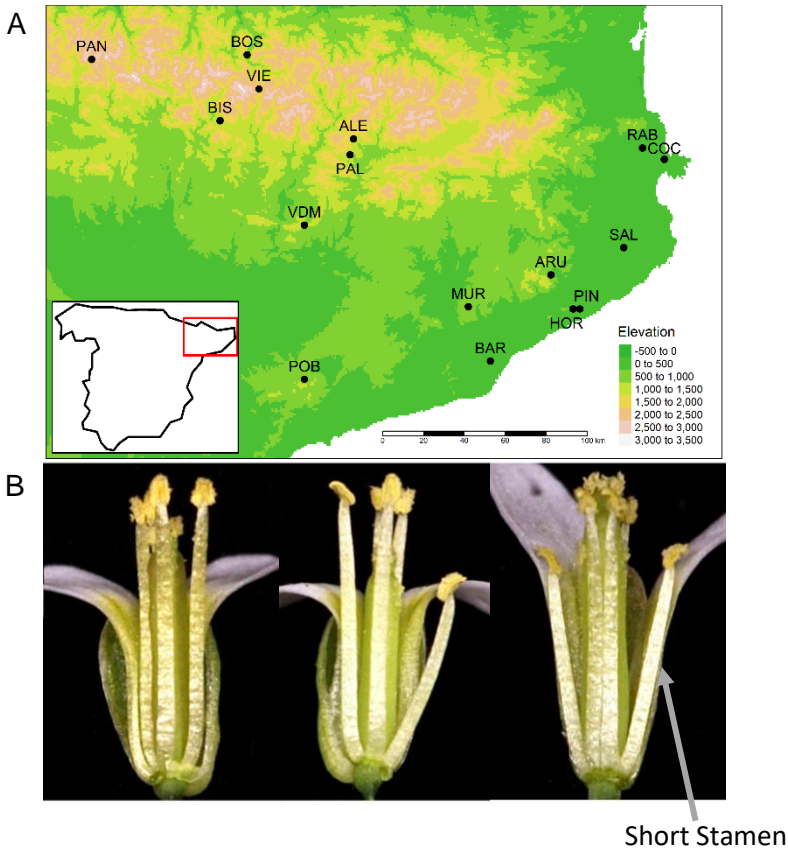
375 Next, we tested if the cline in short stamen loss was maintained by local adaptation using Qpc, an
376 extension of Qst-Fst (Josephs et al., 2019). We found no evidence of local adaptation maintaining the
377 cline in short stamen number (Fig. 3C). One caveat to our findings is that Qst-Fst struggles to identify
378 selection when there is large genetic differentiation between populations or weak selection that results
379 in similar values for Qst and Fst (Whitlock & Guillaume, 2009). This is also potentially an issue for Qpc
380 in this study as there is a great deal of genetic differentiation between high and low elevation populations
381 (Fig. 3A). However, the Qst-Fst approach has identified local adaptation contributing to elevational
382 clines in leaf succulence and specific leaf area in 14 populations of *A. thaliana* from the Swiss Alps where
383 there is also a great deal of genetic differentiation (Luo et al., 2015). Qpc has also previously been used
384 to identify local adaptation in 249 natural *A. thaliana* accessions from across the European native range;
385 local adaptation was identified in initial size, growth rate at 16°C, and temperature response (Clauw et
386 al., 2022). Ultimately, reciprocal transplant studies across elevations in the field will provide the
387 strongest evidence for or against local adaptation.

388 Finally, we used GWAS to identify loci associated with short stamen number. We identified 20 SNPs
389 associated with short stamen number in multiple GWAS (FDR < 0.10). These SNPs lie outside the QTL for
390 short stamens that were previously identified in a cross between a northern European and southern
391 European accession (Royer et al., 2016). The effect sizes of the Royer et al. (2016) QTL (range from 0.05
392 to 0.15) and our loci (Table S4) are comparable, suggesting that we would be able to identify these QTL
393 if they segregated at intermediate frequency in the populations used here. Instead, the populations
394 studied here may be fixed for the southern allele identified in Royer et al (2016). However, our GWAS
395 effect sizes may be inflated due to winner's curse (Göring et al., 2001; Josephs et al., 2017). In addition,
396 the QTL identified in Royer et al. (2016) were epistatic, which could make them harder to detect through
397 standard GWAS approaches. Overall, our results suggest that the latitudinal cline in short stamen
398 number across Europe has a different genetic basis than the elevational cline in short stamen number
399 observed in the Pyrenees.

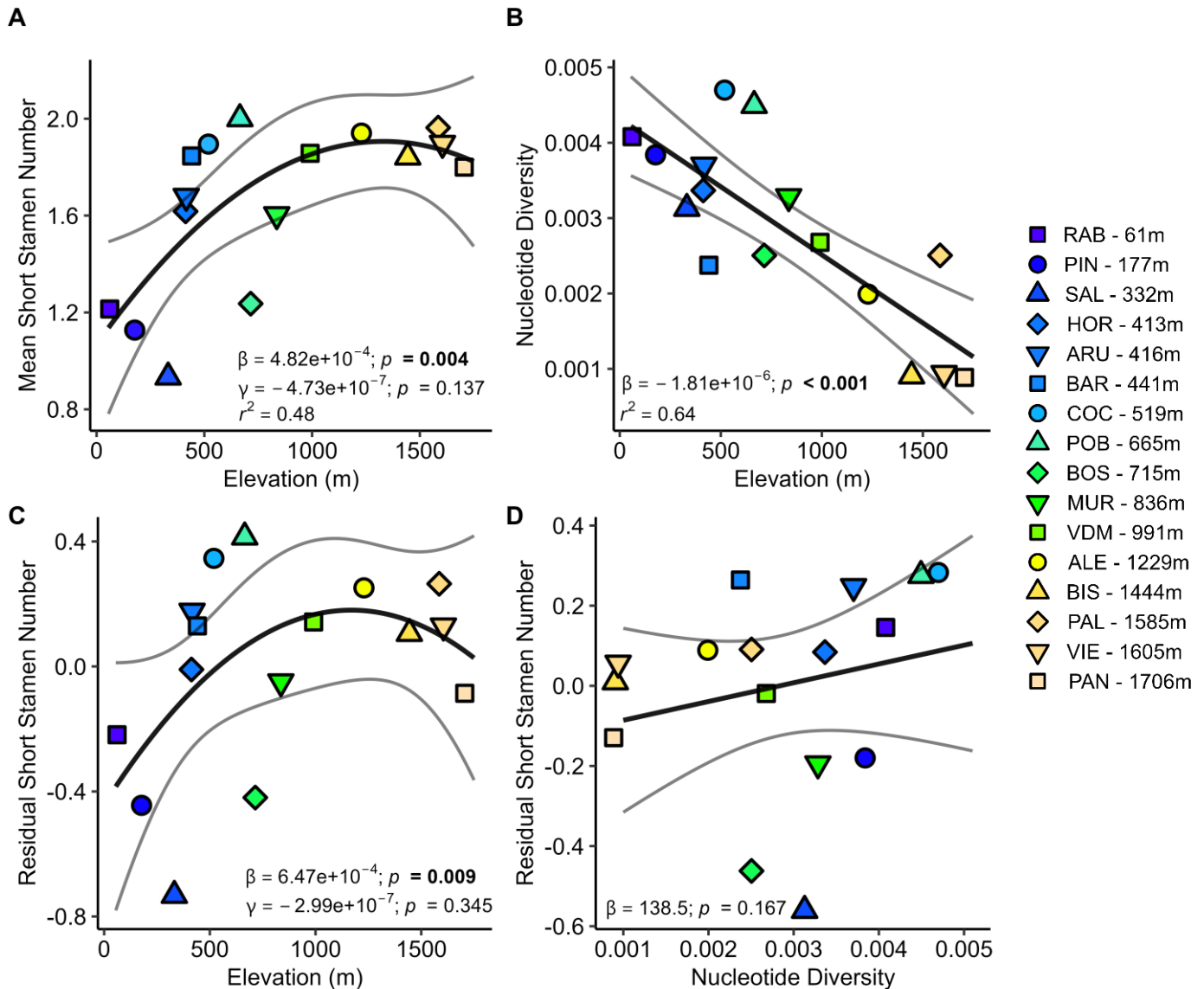
400 While the short stamen loss clines are similar between latitude and elevation, the lack of overlap
401 between the associated loci suggests that short stamen loss is caused by mutations at different genes in
402 different geographic regions. These results are consistent with other findings that the chromosome
403 regions underlying the same phenotypic cline can differ by region. For example, elevational clines of *A.*
404 *thaliana* in different geographic regions show delayed flowering at higher elevations but the genetic
405 basis of this cline varies by global region (Gamba et al., 2022). Additionally, parallel clines in flowering
406 time in *A. thaliana* in the Cape Verde Islands arose from mutations at different genes (FRI and FLC)
407 (Fulgione et al., 2022).

408 In conclusion, short stamen loss is occurring in low elevation populations more than in high elevation
409 populations. Our results suggest that retention of two short stamens in high elevation populations are
410 not explained by reduced effective population sizes or by local adaptation to different optima in high
411 and low elevation populations. Thus, the evolutionary mechanisms underlying variation in short stamen
412 number are excitingly complex and deserve further study. A third hypothesis is that correlated
413 responses to selection may maintain the cline in short stamen number (Futuyma, 2010; Lande, 1979).
414 However, correlated responses to selection on other locally adapted traits could result in weak evidence
415 for divergent selection; we did not find this. Future work should better characterize direct selection and
416 correlated responses to selection on short stamens in the field by measuring fitness of plants with
417 natural variation in short stamen number and experimental manipulation of stamens, like that done by
418 Royer et al. (2016), in the field. Further, refining the short stamen number QTL to candidate genes and
419 comparing to genetic regions underlying locally adapted traits could uncover the role of linked or
420 pleiotropic genes leading to correlated responses to selection. These additional studies across both
421 latitudinal and elevational gradients will characterize the interplay of direct selection, correlated
422 responses to selection, and genetic drift in trait loss and identify parallel evolutionary forces at play
423 across environmental contexts.

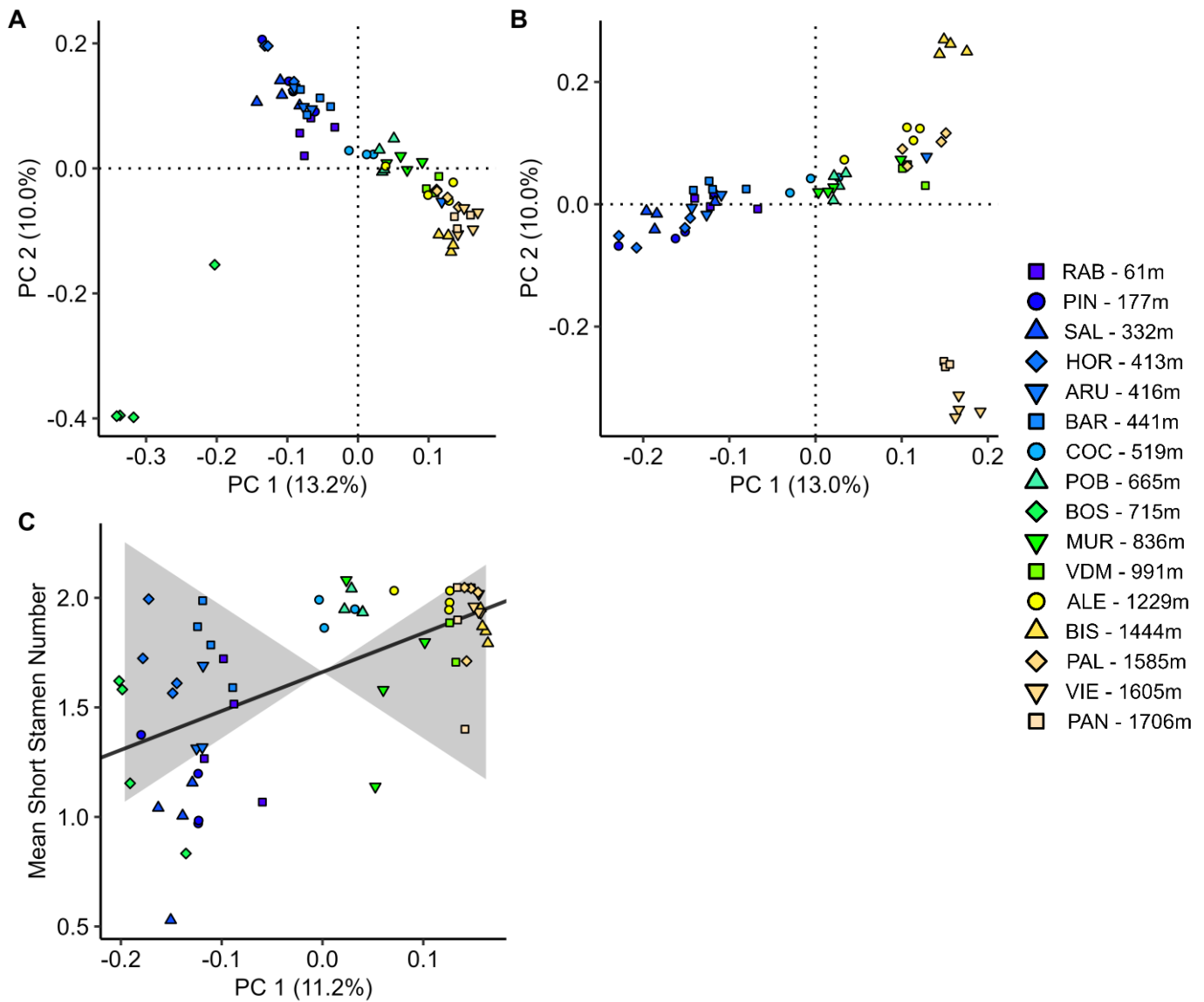
424 **Figures:**



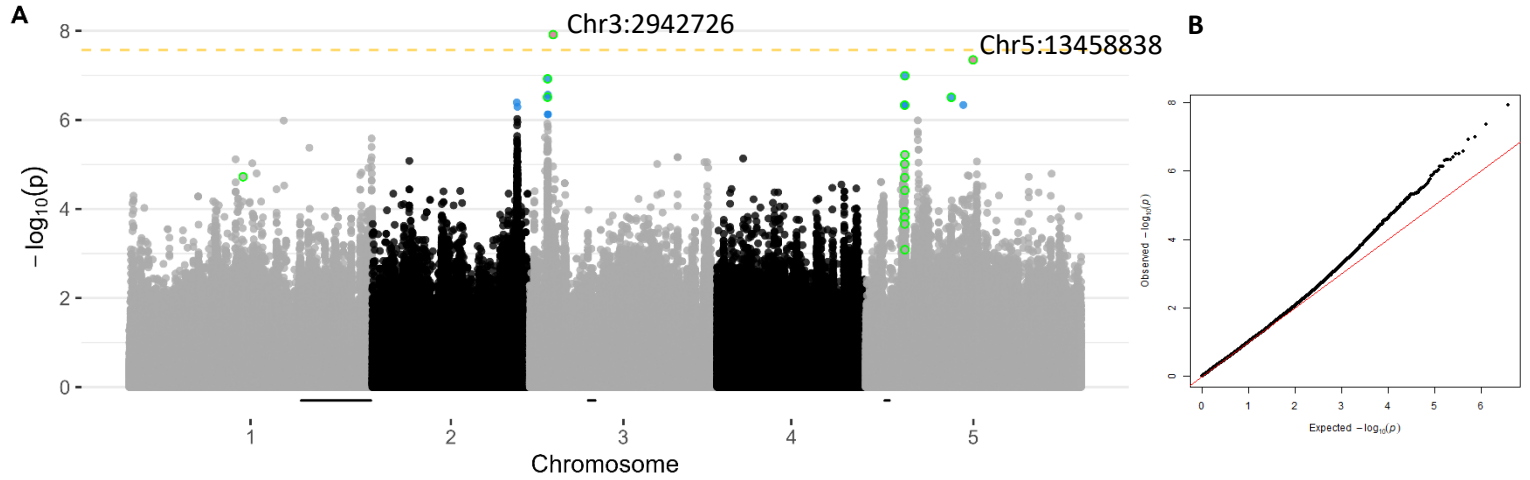
425 **Figure 1: Short stamen number increases with elevation. A)** Each point represents the seed collection
426 site for 16 populations of *A. thaliana* across the Spanish Pyrenees. The map coloring indicates elevation
427 from green at low elevation to white at high elevation. **B)** Natural variation in short stamen number.
428 Flowers have 0 (left), 1 (center), or 2 (right) short stamens. Photos by Frances Whalen.



429 **Figure 2: Effective population size does not explain retention of short stamens at high elevation. A)**
 430 Mean short stamen number shows a quadratic elevational cline. **B)** Elevation strongly predicts
 431 population mean pairwise nucleotide diversity, our measure of effective population size. **C)** The
 432 residuals of the model of short stamen number regressed on nucleotide diversity then regressed on
 433 elevation and **D)** the residuals of the model of short stamen number regressed on quadratic elevation
 434 then regressed on nucleotide diversity. In C and D, statistics from the full model ($r^2 = 0.50$; Table S3) are
 435 displayed on the figures. In all panels, the color of each point represents the population elevation, black
 436 lines are the regression, and grey lines are 95% confidence intervals.



437 **Figure 3: Genetic variation is correlated with elevation, but there is no evidence for divergent**
438 **selection on short stamen number. A)** Genetic PCA for PC1 and PC2 from 1,858,706 SNPs. **B)** Genetic
439 PCA for PC1 and PC2 with BOS individuals excluded. **C)** Qpc results showing the relationship between
440 mean short stamen number and genetic differentiation (black line, $p=0.26$) is within expectations due to
441 neutral evolution (grey shading). The fill color of each point represents the population elevation. Panels
442 A and C used slightly different input data (see Methods).



443 **Figure 4. Few SNPs are associated with short stamen number.** Manhattan plot (A) and QQ plot (B) for
444 arcsine-transformed mean short stamen number. The yellow dashed line indicates significance at $p=0.05$
445 after Bonferroni correction. Blue points are significant below a FDR of 0.10. Pink labelled points are
446 significant below a FDR of 0.05. Points with a green outline are shared between at least two short
447 stamen GWAS below FDR 0.10 ($n = 20$). The black bars on the x axis are Bayes 95% credible intervals for
448 short stamen number QTL identified by Royer et al. (2016).

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